I claim:

- 1. A cDNA encoding for human receptor protein H4-1BB.
- 5 2. The cDNA of claim 1 having a nucleotide sequence as shown in Figure 2.
- 3. The cDNA of claim 1, identified as pH4-1BB deposited at the Agricultural Research Service Culture Collection with the accession number NRRL B21131.
- 4. The cDNA of claim 2 and fragments and derivatives thereof, wherein said fragments and derivatives can be used as a probe to isolate DNA sequences encoding for proteins similar to the receptor protein encoded by said cDNA.

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The receptor protein #4-1BB produced by

- a) inserting the cONA of H4-1BB into an appropriate expression vector,
- b) transfecting said expression vector into an appropriate transfection host,
- c) growing said transfected hosts in appropriate culture media and .
- d) purifying the receptor protein from said culture media.

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6. A protein having the mino acid sequence shown in Figure 2.

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- 30 7. The protein of claim 6 and fragments and derivatives thereof, wherein said fragments and derivatives:
 - a) can be used as a probe to identify ligands to receptor protein H4-1BB;
 - b) can be used to stimulate proliferation B-cell's expressing H4-1BB ligands; or
 - c) can be used to block H4-1RB ligand binding.

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- 8. A monoclonal antibody against H4-1BB which specifically recognizes receptor protein H4-1BB.
- 9. A hybridoma capable of producing a monoclonal antibody 5 against H4-1BB which specifically recognizes receptor protein H4-1BB.
- 10. The method of using the monoclonal antibody of claim 8 to enhance T-cell proliferation comprising the step of 10 treating T-cells that have expressed receptor protein H4-1BB with said monoclonal antibody.
- 11. The method of claim 12 further comprising the step of conducting said treatment in the presence of protein tyrosinase kinase.
- 12. The method of using the monoclonal antibody of claim 8 to enhance T-cell activation comprising the step of treating T-cells that have expressed receptor protein H4-20 1BB with said monoclonal antibody.
 - 13. The method of claim 12 further comprising the step of conducting said treatment in the presence of protein tyrosinase kinase.

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- 14. A fusion protein for detecting cell membrane ligands to human receptor protein H4-1BB, comprising:
 - a) at least a portion of said receptor protein H4-1BB corresponding to the extracellular portion of said receptor protein H4-1BB such that said portion of said receptor protein H4-1BB binds to said cell membrane ligands; and
- b) a detection protein bound to said portion of said receptor protein H4-1BB such that ligand binding can be detected by relative activity assays for said detection protein.

- 15. The fusion protein of claim 14 wherein said detection protein is alkaline phosphatase.
- 16. A method of detecting cell membrane ligands to human 5 receptor protein H4-1BB, comprising:
 - a) providing a fusion protein including:
 - 1) at least a portion of said receptor protein H4-1BB corresponding to the extracellular portion of said receptor protein H4-1BB such that said portion of said receptor protein H4-1BB binds to said cell membrane ligands, and
 - 2) a detection protein bound to said portion of said receptor protein H4-1BB such that ligand binding can be detected by relative activity assays for said detection protein;
 - b) placing said fusion protein in the presence of a cell suspected to express said receptor protein H4-1BB;
 - c) washing said cell of any fusion protein not bound to said cell membrane ligands;
 - d) placing said washed cells in the presence of a substrate for said detection protein and measuring the relative activity of said detection protein.
- 25 17. The method of claim 16 wherein said detection protein is alkaline phosphatase.
- 18. A method of inducing B-cell proliferation comprising the step of treating B-cells that have expressed a ligand 30 to human receptor protein H4-1BB with cells that have expressed receptor protein H4-1BB.

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Math

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